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NEWS 9 Jun 28 ANTE, AQUALINE, BIOENG, CIVILENG, ENVIROENG, MECHENG,
                and WATER from CSA now available on STN(R)
NEWS 10 Jul 12 BEILSTEIN enhanced with new display and select options,
                resulting in a closer connection to BABS
NEWS EXPRESS MARCH 31 CURRENT WINDOWS VERSION IS V7.00A, CURRENT
             MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
             AND CURRENT DISCOVER FILE IS DATED 26 APRIL 2004
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FULL ESTIMATED COST

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10/648,709
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STRUCTURE FILE UPDATES: 20 JUL 2004 HIGHEST RN 713489-00-0 DICTIONARY FILE UPDATES: 20 JUL 2004 HIGHEST RN 713489-00-0

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2004

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Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/reqistryss.html

```
=> s 1,1,13-tetrachloropropane/cn
L1 0 1,1,13-TETRACHLOROPROPANE/CN
```

=> s 1,1,1,3-tetrachloropropane/cn L2 1 1,1,1,3-TETRACHLOROPROPANE/CN

=> d

L2 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2004 ACS on STN

RN 1070-78-6 REGISTRY

CN Propane, 1,1,1,3-tetrachloro- (6CI, 8CI, 9CI) (CA INDEX NAME) OTHER NAMES:

CN 1,1,1,3-Tetrachloropropane

FS 3D CONCORD

MF C3 H4 Cl4

CI COM

LC STN Files: BEILSTEIN*, BIOSIS, CA, CAOLD, CAPLUS, CASREACT, CHEMCATS, CHEMINFORMRX, CHEMLIST, CSCHEM, DETHERM*, GMELIN*, HODOC*, IFICDB, IFIPAT, IFIUDB, MEDLINE, NIOSHTIC, RTECS*, SPECINFO, TOXCENTER, USPATFULL

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DT.CA Caplus document type: Conference; Journal; Patent

RL.P Roles from patents: PREP (Preparation); RACT (Reactant or reagent); USES (Uses); NORL (No role in record)

RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses); NORL (No role in record)

 $C1_3C-CH_2-CH_2C1$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

216 REFERENCES IN FILE CA (1907 TO DATE)

216 REFERENCES IN FILE CAPLUS (1907 TO DATE)

20 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

10/648,709

FILE 'ENCOMPLIT' ACCESS NOT AUTHORIZED

FILE 'ENCOMPLIT2' ACCESS NOT AUTHORIZED

FILE 'ENCOMPPAT' ACCESS NOT AUTHORIZED

FILE 'ENCOMPPAT2' ACCESS NOT AUTHORIZED

COST IN U.S. DOLLARS

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1.5

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=> s 1070-78-6
  48 FILES SEARCHED...
  65 FILES SEARCHED...
           264 1070-78-6
=> s 13 and (iron or ferrous or ferric)
  31 FILES SEARCHED...
  57 FILES SEARCHED...
            48 L3 AND (IRON OR FERROUS OR FERRIC)
=> dup rem 14
DUPLICATE IS NOT AVAILABLE IN 'AQUIRE, BIOCOMMERCE, CAOLD, FEDRIP, GENBANK,
INVESTEXT, KOSMET, RDISCLOSURE, STANDARDS, USAN, DGENE, DPCI, LITALERT,
PCTGEN, PROUSDDR, SYNTHLINE'.
ANSWERS FROM THESE FILES WILL BE CONSIDERED UNIQUE
PROCESSING COMPLETED FOR L4
             38 DUP REM L4 (10 DUPLICATES REMOVED)
=> d 1-38 bib ab
    ANSWER 1 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 1
1.5
AN
    1994:77016 CAPLUS
DN
    120:77016
    Preparation of 1,1-dichloro-3-phenylpropene
TT
IN
    Sagatelyan, Shavarsh A.; Evdokimova, Olga I.
PA
    All-Union Scientific-Research Institute of Chemical Reagents and Pure
     Chemical Substances, Erevan, USSR
SO
    U.S.S.R.
     From: Izobreteniya 1992, (15), 96.
     CODEN: URXXAF
     Patent
DT
LΑ
    Russian
FAN.CNT 1
     PATENT NO.
                 KIND DATE
                                         APPLICATION NO. DATE
                     ----
                           -----
                                          _____
PΙ
    SU 1728213
                      A1
                           19920423
                                          SU 1990-4819244 19900424
PRAI SU 1990-4819244
                           19900424
OS
    CASREACT 120:77016
AB
    The title compound is prepared in a simplified, higher-yielding process by
    alkylation of benzene with 1,1,1,3-tetrachloropropane at reflux in the
    presence of a catalyst comprising a mixture of zinc chloride and
     ferric chloride in a mass ratio of 6:1; the process is conducted
     at a molar ratio of 1,1,1,3-tetrachloropropane to zinc chloride to
     ferric chloride of 1:0.02:0.003.
L5
    ANSWER 2 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
AN
    1991:228314 CAPLUS
DN
    114:228314
ΤI
    Metal complex catalyst for addition reaction of carbon tetrachloride to
     ethylene
ΑU
    Lavrent'eva, E. A.; Ponomarev, V. I.; Lavrent'ev, I. P.
CS
    Inst. Strukt. Makrokin., Chernogolovka, USSR
     Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1990), (12), 2794-8
     CODEN: IASKA6; ISSN: 0002-3353
DT
    Journal
LA
    Russian
    FeIIFeIII2Cl8(AcNMe2)6 catalyzed the title reaction to give ClCH2CH2CCl3
    with 98% selectivity at 55-60% conversion of CCl4. A coordination-ionic
    mechanism was proposed.
```

ANSWER 3 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN

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AN 1989:195133 CAPLUS
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- DN 110:195133
- TI Preparation of catalysts for converting 1,1,1,3-tetrachloropropane to 3,3,3-trifluoropropene by reaction with hydrogen fluoride
- IN Baizer, William X.; Bixler, Jr Robert L.; Meddaugh, Michael D.; Wright, Antony P.
- PA Dow Corning Corp., USA
- SO U.S., 8 pp. CODEN: USXXAM
- DT Patent
- LA English
- FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	US 4798818	Α	19890117	US 1987-125901	19871127
	CA 1326480	A1	19940125	CA 1988-578196	19880926
	JP 01168347	A2	19890703	JP 1988-282627	19881110
	JP 07047127	B4	19950524		
	EP 319153	A1	19890607	EP 1988-310709	19881114
	EP 319153	B1	19920701		
	R: DE, FR,	GB, IT			
DDAT	TTC 1007 100001		10071107		

PRAI US 1987-125901

19871127

OS CASREACT 110:195133

- AB The catalysts with lasting activity are prepared by (A) mech. mixing, in the absence of water, of Al fluoride prepared at <400° and a transition metal compound of Co, Cr, Fe, Mn, Ni, Ti, or V; and (B) contacting the resulting mixture with sufficient anhydrous HF to convert the transition metal compound to a transition metal fluoride. Thus, slowly adding 200 g Al (OH)3 to 400 g 47-52% HF in 1600 mL H2O, maintaining the temperature at 60°, keeping at 60° overnight and drying gave AlF3, which was mixed with 3.2 % FeCl3, heated to 250°, treated with HF for 1 h, heated to 400° with continuous feed of HF for another hour, and cooled under N to 250° to obtain the desired catalyst. The catalyst when used for the title reaction lasted for 34.5 h before losing >10% of its activity.
- L5 ANSWER 4 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 2
- AN 1988:166926 CAPLUS
- DN 108:166926
- TI Reaction of chloroform with ethylene in the presence of pentacarbonyliron-nucleophilic coinitiator systems
- AU Kruglova, N. V.; Petrova, N. A.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Zhurnal Prikladnoi Khimii (Sankt-Peterburg, Russian Federation) (1987), 60(7), 1600-4
 - CODEN: ZPKHAB; ISSN: 0044-4618
- DT Journal
- LA Russian
- OS CASREACT 108:166926
- AB The title telomerization with excess Me2CHOH, HMPA or DMF as the nucleophile and 2:1 CHCl3-C2H4 gave Cl2CHCH2CH2Cl as the major product, along with 2-9% ClCH(CH2)4Cl, 3-5% CCl2(CH2CH2Cl)2, and, with small nucleophile excesses, lesser amts. of telomers resulting from initial C-H cleavage. The catalyst selectivity increased in the stated order of nucleophiles.
- L5 ANSWER 5 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 3
- AN 1988:186812 CAPLUS
- DN 108:186812
- TI Addition of 1,1,1-trichloroalkanes to trimethylvinylsilane in the presence of systems based on **iron** pentacarbonyl
- AU Kamyshova, A. A.; Dostovalova, V. I.; Chukovskaya, E. Ts.

- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1987), (5), 1174-7 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- OS CASREACT 108:186812
- AB Addition reaction of RCCl3 (R = Me, CH2ClCH2) with CH2:CHSiMe3 in the presence of Fe(CO)5 and Ph3P, DMF, HMPA or Me2CHOH gave RCCl2CH2CHClSiMe3 (I). Dehydrochlorination of I (R = CH2ClCH2) with Fe(CO)5-PPh3 gave 71% of a mixture of (Z)-clcH2CH:CClCH2CHClSiMe3 and ClCH2CH2CCl:CHCHClSiMe3.
- L5 ANSWER 6 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 4
- AN 1986:555080 CAPLUS
- DN 105:155080
- TI 1,1,1,3-Tetrachloropropane
- IN Astrologes, Gary W.
- PA Halocarbon Products Corp., USA
- SO U.S., 5 pp.
- CODEN: USXXAM
- DT Patent
- LA English
- FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	US 4605802	Α	19860812	US 1982-364175	19820401
	CA 1273645	A1	19900904	CA 1986-506579	19860414
	JP 62263134	A2	19871116	JP 1986-104996	19860509
	EP 247215	A1	19871202	EP 1986-107098	19860526
	R: CH, DE,	FR, GB	, IT, LI, NL		
PRAI	US 1982-364175		19820401		

- OS CASREACT 105:155080
- Ethylene is reacted with CCl4 in the presence of a phosphite ester [especially P(OEt)3 or P(OBu)3] and powdered Fe catalyst to yield 1,1,1,3-tetrachloropropane (I). Optionally, added FeCl3 may be present. The powdered Fe is especially electrolytic Fe powder or H-reduced Fe powder. In an autoclave at 90-110 psig C2H4, CCl4 .apprx.900, P(OEt)3 5.5, electrolytic Fe powder 7.5, and FeCl3 1.174 g were reacted at 95°, resulting in 75.9% I and 21.2% unreacted CCl4 in the product, corresponding to 96.3% yield of I.
- L5 ANSWER 7 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 5
- AN 1987:195822 CAPLUS
- DN 106:195822
- TI Molybdenum hexacarbonyl as a catalyst for dehydrochlorination of compounds containing a trichloromethyl group
- AU Kuz'mina, N. A.; Blinova, N. V.; Chukovskaya, E. Ts.; Grechkina, E. M.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1986), (5), 1139-42 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- OS CASREACT 106:195822
- AB Heating CCl3CH2R [I; R = n-hexyl, (CH2)3Cl, CH2Cl, CHClC5H11-n] with 0.03 equiv Mo(CO)6 gave 14-98% CCl2:CHR (II; same R) and 0-10% Cl2CHCH2R (same R) with 70-100% I conversion. I (R = n-hexyl) gave 74% II with 0.03 equiv 1:1 Fe(CO)5-Ph3P, but only 30% II with no Ph3P. Other M(CO)6 (M = W, Cr) or Mn2(CO)10 were ineffective catalysts alone or with Ph3P or DMF.
- L5 ANSWER 8 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 6
- AN 1987:195821 CAPLUS
- DN 106:195821

- TI Hydrogenolysis of a carbon-chlorine bond in a dichloromethylene group in polychloroalkanes initiated by pentacarbonyliron, decacarbonylmanganese, or tert-butyl peroxide with triethylsilane
- AU Kiseleva, L. N.; Rybakova, N. A.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1986), (5), 1136-8 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- OS CASREACT 106:195821
- AB Reduction of ClCH2CH2CCl2CH2CHClR (I; R = H, Bu, n-pentyl) with 1 equiv Et3SiH and 5-20% of the title initiators gave 71-92% ClCH2(CH2CHCl)2R (II; same R) with 35-96% I conversion. Using 20% Mn2(CO)10 gave 85-90% II with 82-96% I conversion.
- L5 ANSWER 9 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 7
- AN 1985:203584 CAPLUS
- DN 102:203584
- TI 1,1,2,3-Tetrachloropropene
- IN Woodard, Scott Santford
- PA Monsanto Co. , USA
- SO Eur. Pat. Appl., 30 pp.
- CODEN: EPXXDW
 DT Patent
- LA English
- FAN.CNT 1

t. Little .	CTA T	_										
	PA	TENT NO.		KI	ND	DATE			API	PLICATION	NO.	DATE
PI	EΡ	131560		Α	1	1985	0116		EP	1984-870	0092	19840704
	EP	131560		В	1	1987	0812					
		R: AT,	BE,	CH,	DE	, FR,	GB,	IT,	LI, 1	LU, NL		
	US	4535194		A		1985	0813		US	1983-513	1131	19830706
	AT	28853		E		1987	0815		AT	1984-870	0092	19840704
	DK	8403297		Α		1985	0107		DK	1984-329	97	19840705
	AU	8430318		Α	1	1985	0110		AU	1984-303	318	19840705
	JΡ	60036429)	Α	2	1985	0225		JP	1984-138	3112	19840705
	JP	02047969)	В	4	1990	1023					
	ZA	8405179		Α		1985	0327		ZA	1984-517	79	19840705
	HU	34422		Α	2	1985	0328		HU	1984-264	48	19840705
	CA	1230132		Α	1	1987	1208		CA	1984-458	3213	19840705
	SU	1452476		Α	3	1989	0115		SU	1984-375	57914	19840705
	US	4650914		Α		1987	0317		US	1985-690	8000	19850109
	SU	1470174		Α	3	1989	0330		SU	1986-400	00868	19860106
	CA	1234156		Α	2	1988	0315		CA	1987-531	1178	19870304
	CA	1234157		A	2	1988	0315		CA	1987-531	1179	19870304
PRAI	US	1983-511	131			1983	0706					
	ΕP	1984-870	092			1984	0704					
	CA	1984-458	213			1984	0705					

- OS CASREACT 102:203584
- AB The FeCl3-catalyzed isomerization of 2,3,3,3-tetrachloropropene (I) gave 1,1,2,3-tetrachloropropene. Thus, 1,1,1,2,3-pentachloropropane was dehydrochlorinated to I, and I was heated with FeCl3 at 103° to give 1,1,2,3-tetrachloropropene.
- L5 ANSWER 10 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1985:453693 CAPLUS
- DN 103:53693
- TI Monoadducts of olefins and telogens reactive therewith
- IN Woodard, Scott Santford
- PA Monsanto Co. , USA
- SO Eur. Pat. Appl., 44 pp. CODEN: EPXXDW

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DT Patent
LA English
FAN.CNT 1
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PA	TENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI EP	131561	A1	19850116	EP 1984-870094	19840704
EP	131561	B1	19861015		
	R: AT, BE,	CH, DE	, FR, GB, IT,	LI, LU, NL	
AT	22877	E	19861115	AT 1984-870094	19840704
DK	8403298	Α	19850107	DK 1984-3298	19840705
AU	8430317	A1	19850110	AU 1984-30317	19840705
JP	60036428	A2	19850225	JP 1984-138113	19840705
ZA	8405182	Α	19850327	ZA 1984-5182	19840705
HU	34421	A2	19850328	HU 1984-2649	19840705
PRAI US	1983-511130		19830706		
EP	1984-870094		19840704		

- AB A monoadduct of a taxogen and a telogen was prepared in the presence of a metallic Fe source and a P(V) promoter containing a phosphoryl group. Thus, CCl4, FeCl3, (EtO)3PO, and Fe wire were pressurized with C2H4 and the mixture stirred at 120° to give 9.6% yield of CCl3CH2CH2Cl. Other P promoters included (BuO)3PO, OP(OEt)2CH:CH2, and OP(OEt)2OH. Mild steel bars and stainless steels were also used in place of Fe wire.
- L5 ANSWER 11 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 8
- AN 1986:148054 CAPLUS
- DN 104:148054
- TI EPR study of ·CCl2CHX(CH2)3R radical rearrangement involving a 1,5-hydrogen shift and transformations of these radicals
- AU Gasanov, R. G.; Vasil'eva, T. T.; Kruglova, N. V.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin. im. Nesmeyanova, Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1985), (11), 2507-12 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- OS CASREACT 104:148054
- AB A spin-trapping study confirmed the rearrangement of RCH2(CH2)3CCl2• (R = Me, Et, MeO) to HCCl2(CH2)3CHR• (same R) via a 1,5-H migration. MeO(CH2)3CHRCCl2• (R = H, CCl3) isomerized to HCCl2CHRCH2CH2CH(OMe)•, which were oxidized to HCCl2CHRCH2CH2CO2Me.
- L5 ANSWER 12 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1983:445947 CAPLUS
- DN 99:45947
- TI EPR study of the interaction of **iron** carbonyl (Fe(CO)5) and metal carbonyl (M(CO)6) with halogen-containing compounds in the presence of triphenylphosphine and M(CO)5PPh3 (M = chromium, molybdenum, tungsten)
- AU Gasanov, R. G.; Ivanova, L. V.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin. im. Nesmeyanova, Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1983), (5), 1045-53 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- AB Photoreaction of MCOn (M = Fe, Cr, Mo, W) with CCl4 in the presence of PPh3 was investigated by ESR. Substitution of CO by PPh3 led to the increase of the yield and activity of the metal-containing transients responsible for the generation of the organic radicals. During photolysis of substituted carbonyls of Cr, Mo and W in general cleavage of the metal-CO and not the metal-PPh3 bond occurred.
- L5 ANSWER 13 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1983:160189 CAPLUS
- DN 98:160189

- TI Synthesis of polychloroalkanes with various chlorine containing groups
- AU Zhiryukhina, N. P.; Kamyshova, A. A.; Chukovskaya, E. Ts.; Freidlina, R.
- CS Inst. Elementoorg. Soedin. im. Nesmeyanova, Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1983), (1), 152-7 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- OS CASREACT 98:160189
- AB Reaction of CH2:CMeCl with CCl3CH2CHCl2 in the presence of Fe(CO)5 and (Me2N)3PO gave 42-60% MeCCl2CH2CCl2CH2CH2Cl along with 1-3% MeCHClCH2CCl2CH2CH2Cl and MeCHClCH2CCl2CH2CHCl2. CH2:CMeCl and CCl3CHCl2 gave 41-53% MeCCl2CH2CCl2CHCl2 and 7-8% MeCCl2CH2CCl:CHCl (Z-E 3.3:1).
- L5 ANSWER 14 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 9
- AN 1982:526953 CAPLUS
- DN 97:126953
- TI Reaction of 1,1,1,3-tetrachloropropane with 1-hexene initiated by pentacarbonyliron, hexacarbonylmolybdenum, and decacarbonyldimanganese
- AU Chukovskaya, E. Ts.; Kuz'mina, N. A.; Rozhkova, M. A.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1982), (6), 1345-9 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- OS CASREACT 97:126953
- AB Fe(CO)5 was the most effective catalyst for the title reaction at 120-40° in the presence of Me2CHOH cocatalyst, but Mn2(CO)10 also catalyzed the reaction in the absence of Me2CHOH. The major product was ClCH2CH2CCl2CH2CHClBu (I), and the I-ClCH2CH2CCl2CH2CH2Bu ratio depended on the catalyst.
- L5 ANSWER 15 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 10
- AN 1980:110458 CAPLUS
- DN 92:110458
- TI Addition of 1,1,1,3-tetrachloropropane to 1-hexene initiated by the iron pentacarbonyl + HMPA + N,N-dichloro-p-chlorobenzenesulfamide system
- AU Rybakova, N. A.; Chukovskaya, E. Ts.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1979), (11), 2618-20 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- OS CASREACT 92:110458
- AB Including p-ClC6H4SO2NCl2 (I) in the title initiator mixture increased the ClCH2CH2CH2CH2CHClBu yield from the title reaction at 105° in air to 96%, vs. 40% in the absence of I.
- L5 ANSWER 16 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1979:574753 CAPLUS
- DN 91:174753
- TI Reduction of polychloroalkanes of CCl3CH2CHXY type initiated by iron pentacarbonyl
- AU Vasil'eva, T. T.; Kruglova, N. V.; Tarasova, T. V.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1979), (7), 1543-8 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- OS CASREACT 91:174753
- AB Fe(CO)5-initiated reduction of CCl3CH2CHXR (X = Cl, R = H, Me; X = H, R = Me)

with Me2CHOH gave Cl2CHCH2CHXR as the major products, along with products derived from recombination of the intermediate \bullet CCl2CH2CHXR (I) followed by dechlorination (e.g., MeCHClCH2CCl:CClCH2CHClMe). No evidence for 1,3-H shift in I was observed

- L5 ANSWER 17 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1979:21926 CAPLUS
- DN 90:21926
- TI Reaction of polychloroalkanes with pentacarbonyliron and hexametapol in the presence or absence of olefins studied by an EPR method using spin traps
- AU Freidlina, R. Kh.; Gasanov, R. G.; Grigor'ev, N. A.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Doklady Akademii Nauk SSSR (1978), 242(2), 354-7 [Chem.] CODEN: DANKAS; ISSN: 0002-3264
- DT Journal
- LA Russian
- AB The yields of C13C•, C1CH2CH2CC12•, and C1CH2CH2CC12CH2CHBu• radicals, formed in the photoaddn. of C1CH2CH2CC13 to 1-hexene with a Fe(CO)5-(Me2N)3PO catalyst, proceeded through sep. maximum at different concns. of catalyst components. This showed that self-inhibition occurred in the initiation stage. The radicals were trapped with nitrosodurene or tert-butylphenylnitrone.
- L5 ANSWER 18 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1978:546097 CAPLUS
- DN 89:146097
- TI Study of the reaction of carbon tetrachloride and 1,1,1,3tetrachloropropane with 1-hexene, initiated by pentacarbonyliron and nucleophilic cocatalysts, by an EPR method using spin traps
- AU Gasanov, R. G.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Doklady Akademii Nauk SSSR (1978), 239(1), 94-7 [Chem.] CODEN: DANKAS; ISSN: 0002-3264
- DT Journal
- LA Russian
- AB The title photoreaction was carried out with Me2CHOH, DMF, or (Me2N)3PO as cocatalyst or without cocatalyst, and the nitroxyl radicals formed with nitrosodurene (I), Me3CNO, or α -phenyl-N-tert-butylnitrone as trapping agent were analyzed by ESR. The radical yields obtained in the 1-hexene-CCl4-Fe(CO)5-I system were higher with cocatalysts than without them, and the increase depended on the nature and concentration of the cocatalyst.
- L5 ANSWER 19 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1977:501917 CAPLUS
- DN 87:101917
- TI Tetrachloroalkanes
- IN Takamizawa, Minoru; Okamoto, Haruo; Umemura, Mitsuo; Kohya, Kazuo
- PA Shin-Etsu Chemical Industry Co., Ltd., Japan
- SO Jpn. Kokai Tokkyo Koho, 4 pp. CODEN: JKXXAF
- DT Patent
- LA Japanese
- FAN.CNT 1

ΡI

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	JP 52059102	A2	19770516	JP 1975-134901	19751110
7 T	TD 1005 104001		4 6 5 5 4 4 4 6		

PRAI JP 1975-134901 19751110

AB CCl3CH2CH2Cl (I), CCl3CH2CH2CH2CH2Cl (II), and CCl3CH2CHClMe were prepared by reaction of C2H4 or propene with CCl4 in the presence of HC(OEt)3, and FeCl2 or FeCl3. Thus, C2H4 was introduced into a mixture of CCl4 800,

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HC(OEt)3 7.7, and FeCl3.6H2O 1.4 g in N until the pressure rose to 5
kg/cm2 gage. The mixture was stirred 6 h at 120° while introducing
C2H4 to keep the pressure at 10 kg/cm2 gage to give 329 g I and 37 g II
(total 93.2% yield).
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- ANSWER 20 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN 1.5
- 1977:170762 CAPLUS AN
- DN 86:170762
- ΤI Reactions of gem-trichloroalkanes with unsaturated compounds in the presence of iron carbonyl (Fe(CO)5) and a cocatalyst
- AII Freidlina, R. Kh.; Kuz'mina, N. A.; Kamyshova, A. A.; Polishchuk, S. D.; Chukovskaya, E. Ts.
- Inst. Elementoorg. Soedin, Moscow, USSR CS
- Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1977), (1), 174-7 SO CODEN: IASKA6; ISSN: 0002-3353
- Journal DT
- Russian LΑ
- Telomerization of CH2:CHCl with RCH2CCl3 (I; R = Me, Cl2CH) in the AB presence of Fe(CO)5 and Me2CHOH at 130-45° in a sealed ampul afforded MeZ(CH2CHCl)nCl (Z = CH2CCl2, CH:CCl; n = 2, 3) and (Cl2CHCH2)2CCl2, resp. Analogous reaction of I (R = ClCH2) with CH2:CHCH2Cl gave ClCH2CH2CCl2CH2R1 (R1 = CHClCH2Cl, CH:CH2) and (ClCH2CH2CCl2CH2)2CHCl. Addition of I (R = H, ClCH2) to CH2:CMe2 at 125-7° in the presence of Fe(CO)5 and (Me2N)3PO yielded RCH2CCl2CH2CMe2Cl and traces of RCH2CCl2CH2CMe:CH2.
- ANSWER 21 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN L5
- ΑN 1975:427447 CAPLUS
- DN 83:27447
- ΤТ Reactions of 1,1,1,3-tetrachloropropane with α -olefins catalyzed by triethyl phosphite-ferrous or ferric chloride
- ΑIJ Sato, Toru; Kurita, Naoyasu; Kise, Hideo; Seno, Manabu; Asahara, Teruzo
- CS Inst. Ind. Sci., Univ. Tokyo, Tokyo, Japan
- Nippon Kagaku Kaishi (1975), (2), 398-9 SO CODEN: NKAKB8; ISSN: 0369-4577
- DТ Journal
- LA Japanese
- The telomerizations of 1,1,1,3-tetrachloropropane (I) with α -olefins AB were carried out in the presence of P(OEt)3-FeCl2 or -FeCl3. The main products were the simple adducts 1-substituted 1,3,3,5tetrachloropentanes. The reaction proceeds by abstraction of Cl from I. The conversion of the olefin to the adduct increased with an increase in I to olefin ratio.
- L5 ANSWER 22 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN1974:551521 CAPLUS
- DN 81:151521
- TI1,1,3-Trichloro-1-propene from 1,1,1,3-tetrachloropropane
- IN Fujimori, Kunihiko; Chiyomaru, Isao; Kubota, Renji; Takita, Kiyoshi
- Kumiai Chemical Industry Co., Ltd. PA
- Jpn. Kokai Tokkyo Koho, 2 pp. SO
- CODEN: JKXXAF
- DT Patent
- LΑ Japanese
- FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	JP 49066613	A2	19740627	JP 1972-107662	19721027
PRAI	JP 1972-107662		19721027		

Cl3CCH2CH2Cl (I) was dehydrochlorinated to Cl2C:CHCH2Cl (II) using anhydrous FeCl3 as catalyst. Thus, I was heated with 0.2-0.6 g FeCl3 4-5 hr at 80-95° to give 85-6% II. The yield was lowered to 20-44% with >1 g

or <0.1 g FeCl3.

- L5 ANSWER 23 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1973:536440 CAPLUS
- DN 79:136440
- TI Telomerization of methyl acrylate and methyl methacrylate by 1,1,1,3-tetrachloropropane initiated by iron pentacarbonyl and a nucleophilic reagent
- AU Chukovskaya, E. Ts.; Rozhkova, M. A.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin, Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1973), (8), 1782-8 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- ClCH2CH2CCl3 in telomerization with MeO2CCH:CH2 and HO2CCH:CH2 with large excess of the telogen and either isoamyl alc. or 2-ethylhexanol as the nucleophilic reagent gave at 110-40° only 3-5% yields of products, but in the presence of PhNMe2 or Me2NCHO both adducts and telomers were formed in 2 series: Cl(CH2)2CCl2(CH2CXCO2Me)nCl and Cl(CH2)2CCl2(CH2CXCO2Me)nH (X = H, Me, n = 1-3). Both are formed by reaction of the same radical intermediate with the Cl donor being the telogen and H donor being the nucleophile. The reactions were run as described earlier (1969) with Fe(CO)5 as the principal initiator aided by the added nucleophile. Yields of products were tabulated for this reaction and for similar ones with CH2:CHCN and CH2:CMeCO2Me. The products were hydrogenated over Pd/BaSO4. Replacement of Cl by H leads to mixed products, but may be a source of Cl(CH2)2CCl2(CH2CXCO2Me)nZ (X = H, Me, Z = H, Cl).
- L5 ANSWER 24 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1973:491497 CAPLUS
- DN 79:91497
- TI Reaction of ethylene with $\alpha, \alpha, \alpha, \beta$ and $\alpha, \alpha, \alpha, \beta, \beta$ -polychloroalkanes initiated by **iron** pentacarbonyl and isopropanol
- AU Karapet'yan, Sh. A.; Belyavskii, A. B.; Kuz'mina, N. A.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1973), (6), 1272-5 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- The title reaction in the presence of varying amounts of FeCl2 hydrate, Fe + HCl, FeCl2-FeCl3, or Fe(CO)5 was studied for CCl3R [R = CHCl2, CH2Cl, (CH2)2Cl, CHCl(CH2)2Cl, CCl2(CH2)3Cl]. Addition of such compds., with 4th Cl in terminal position, to C2H4 occurred readily in iso-PrOH in the presence of either Fe(CO)5 alone or mixed with Fe chlorides or by the action of a steel plate and a small amount of HCl. Similarly facile was the reaction of 1,1,1-trichloroalkanes. In 1,1,1,2- or 1,1,1,2,2-polychloroalkanes, however, dechlorination took place that led to di- and trichloroalkenes, resp. In the ease of such loss of Cl the decreasing tendency follows the series CCl3CCl2R, CCl3CHClR, CCl3CH2Cl, and CCl3CHCl2. The ease of addition to C2H4 is the opposite of the above order.
- L5 ANSWER 25 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1973:123925 CAPLUS
- DN 78:123925
- TI Reaction of thiols with 1-heptene and 1,1,1,3-tetrachloropropane in the presence of **iron** pentacarbonyl
- AU Kandror, I. I.; Petrova, R. G.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1973), (1), 167-70 CODEN: IASKA6; ISSN: 0002-3353

- DT Journal
- LA Russian
- AB The yields of products formed by the title reaction of PhSH and BuSH in the presence or absence of Fe(CO)5, FeCl3, or azobisisobutyronitrile were reported for reactions run at 75° or 140°. At the lower temperature the tetrachloride acts as inert diluent only and the RSH addition to
 - the olefin is wholly inhibited by Fe(CO)5. At 140° the thiols reduce the tetrachloride to 1,1,3-trichloropropane and the effectiveness of Fe(CO)5 as an inhibitor of radical addition of RSH to olefins drops.
- L5 ANSWER 26 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1973:3603 CAPLUS
- DN 78:3603
- TI Addition of 1,1,1-trichloroethane and 1,1,1,3-tetrachloropropane to substituted olefins in the presence of iron pentacarbonyl and a nucleophilic additive
- AU Chukovskaya, E. Ts.; Kuz'mina, N. A.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1972), (9), 2110-12 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- AB Slow addition of Me2NPh, Fe(CO)5, CH2:CHOAc, and CCl3CH2CH2Cl (I) over 3.5 hr to an equimolar amount I at 120-40°, then heating 1 hr at 120° gave 1,3,3,5-tetrachloro-1-acetoxypentane. CH2:CHOAc and CCl3Me gave 15% 1.3,3-trichloro-1-acetoxybutane. Chloroprene similarly gave a mixture of 70% 1.4- and 30% 4,1-adducts [Cl(CH2)2CCl2CH2CCl:CHCH2Cl and Cl(CH2)2CCl2CH2CH:CClCH2Cl], each of which existed in Z- and E-forms, with the Z-isomer predominant.
- L5 ANSWER 27 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1972:487798 CAPLUS
- DN 77:87798
- TI Addition of 1,1,1,3-tetrachloropropane to acrylonitrile in the presence of iron pentacarbonyl and nucleophilic additives
- AU Kuz'mina, N. A.; Chukovskaya, E. Ts.; Rozhkova, M. A.; Freidlina, R. Kh.
- CS Inst. Elementoorg. Soedin., Moscow, USSR
- SO Izvestiya Akademii Nauk SSSR, Seriya Khimicheskaya (1972), (4), 961-3 CODEN: IASKA6; ISSN: 0002-3353
- DT Journal
- LA Russian
- AB Addition of CCl3CH2CH2Cl to CH2:CHCN in the presence of Fe(CO)5 gave various proportions of ClCH2CH2CCl2CH2CHClCN (I) and ClCH2CH2CCl2CH2CH2CN (II). In the presence of Me2NCHO I is favored, and PhNMe2 favors II. Alcs. or MeCN lead to a mixture of I and II in low yields (4-6%), whereas with the best solvents the total yields were 31-5%. The reactions were run at 130-40° for .apprx.0.5 hr.
- L5 ANSWER 28 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1972:527093 CAPLUS
- DN 77:127093
- TI Reaction of ethylene with 1,1,1,3-tetrachloropropane initiated by hexaethylphosphorous-triamide-ferric chloride
- AU Sato, Toru; Kurita, Ariyasu; Seno, Manabu; Asahara, Teruzo
- CS Inst. Ind. Sci., Tokyo Univ., Tokyo, Japan
- SO Yuki Gosei Kagaku Kyokaishi (1972), 30(5), 462-4 CODEN: YGKKAE; ISSN: 0037-9980
- DT Journal
- LA Japanese
- AB The telomerization reaction of ethylene [74-85-1] with 1,1,1,3-tetrachloropropane [1070-78-6] initiated by

hexaethylphosphorous triamide [2283-11-6]-iron(III) chloride [7705-08-0] was investigated in a stainless steel autoclave under various reaction conditions (by changing temperature, ethylene pressure and composition of

initiator). The reaction products were α , γ , γ , ω -tetrachloroalkane homologs which were the same as those from the reactions initiated by triethyl phosphite [122-52-1]-ferric salts; 80-90 weight % of the products were 1:1 telomer, 1,3,3,5-tetrachloropentane [24616-07-7].

- L5 ANSWER 29 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1971:448334 CAPLUS
- DN 75:48334
- TI Telomerization of ethylene and carbon tetrachloride
- AU Asahara, Teruzo; Sato, Toru
- CS Inst. Ind. Sci., Univ. Tokyo, Tokyo, Japan
- SO Kogyo Kagaku Zasshi (1971), 74(4), 703-5 CODEN: KGKZA7; ISSN: 0368-5462
- DT Journal
- LA Japanese
- AB The telomerization of ethylene and carbon tetrachloride at 160°/60-70 atm in the presence of Et3PO3-metal salt gave the highest yield of carbon tetrachloride-ethylene telomer when FeCl2 or FeCl3 was used as the metal salt. The telomer obtained in the highest yield contained >90% 1,1,1,3-tetrachloropropane. A free radical mechanism was discussed.
- L5 ANSWER 30 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1969:412514 CAPLUS
- DN 71:12514
- TI Adducts of carbon tetrachloride and ethylene, propylene, or butadiene
- IN Asscher, Meir; Katchalsky, Aharon; Vofsi, David
- SO Brit., 6 pp. Addn. to Brit. 1023423
- CODEN: BRXXAA
- DT Patent
- LA English
- FAN.CNT 1

PΙ

CIVI				
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
GB 1146463		19690326	GB	19660322

AB Cu or Fe salts catalyze the homogeneous reaction at elevated temperature of CC14

with ethylene, propylene, or butadiene. Thus, 15.4 g. CCl4, 2 g. iso-PrOH, 356 mg. ferric acetylacetonate, and 212 mg. benzoin in a 200-ml. Ag-lined autoclave were treated with 14 g. C2H4 at 100° and 1200 psi. for 12 hrs. to give 11 g. product, b25 60-140°, 55:45 ClCH2CH2CCl3ClCH2CH2CH2CH2CCl3; without benzoin, the yield was 1.5 g. Similarly, with CCl4, CH2Cl2, Fe naphthenate, NEt3.HCl, and benzoin, C3H6 yielded 50% MeCHClCH2CCl3, or without NEt3-HCl, 38%; with CCl4, MeCN, and cupric acetylacetonate, 5.4 g. C4H6 yielded 6.5 g. ClCH2CH:CHCH2CCl3, b25 76-150°, and 2.6 g., b15 85-160°, mainly of mixed isomeric tetrachlorononadienes. The use of cupric stearate with ethylene or butadiene is disclaimed.

- L5 ANSWER 31 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 1967:28319 CAPLUS
- DN 66:28319
- TI Polymerization of ethylene. IV. The reaction between ethylene and carbon tetrachloride initiated by a redox catalyst systems
- AU Mogi, Noboru; Takahama, Hiroshi; Takahashi, Akira
- CS Showa Denko Co., Tokyo, Japan
- SO Kogyo Kagaku Zasshi (1966), 69(6), 1218-24

10/648,709 CODEN: KGKZA7; ISSN: 0368-5462 Journal DT LΑ Japanese cf. CA 61, 4196h; 63, 16201c. The reaction between C2H4 and CCl4 was AΒ initiated efficiently by organic peroxide-reducing agent-metal carboxylate catalyst systems at relatively low temperature The reducing agent was benzoin and an amine such as aminoethylethanolamine or mono-, di-, or triethanolamine. For example, 1 + 10-3, 1 + 10-3, and 1 + 10-4 mole, resp., of Bz2O2, (I) benzoin, and metal (Fe, Pb, Zn, Co, Ni, or Mn) naphthenate were added to 1 mole of CC14, and C2H4 was bubbled into the reaction mixture at 50° for several hrs. The reaction rate was estimated from the rate of C2H4 consumption measured with a gas buret. The reaction products were analyzed by gas chromatography and substantially no compds. other than 1,1,1,3-tetrachloropropane were detected. Among the metal naphthenates tested, Fe naphthenate (II) was by far the most effective, while the kinds of acid groups of the Fe carboxylates had little effect on the effectiveness of the catalyst systems. The effectiveness of different kinds of organic peroxides was tested and it was found that lauroyl peroxide was the most effective, while cumene hydroperoxide was completely ineffective in the organic peroxides-benzoin-II catalyst systems. The reaction rate was proportional to the concentration of the organic peroxide, e.g. dicumyl peroxide (III), in the concentration range (III/CCl4 mole ratio) 1.0-2.0 + 10-3 with the ratios benzoin/III = 1:1 and II/III = 0.1:1 and at 50°. The effects of the concns. of benzoin and II were also investigated. The apparent activation energy for the reaction calculated from the temperature dependence reaction rate was 6.4 or 8.5 kcal./mole when III or I, resp., was combined with benzoin and II. The value for the catalyst system I-triethanolamine-II was 7.5 kcal./mole. A considerable reaction rate was obtained even in the absence of I or II from the I-triethanolamine-II system and there seems to be an additivity between the reaction rates with both binary catalyst systems that gives the rate with the ternary catalyst system. ANSWER 32 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN L5 1964:22978 CAPLUS AN 60:22978 DNOREF 60:4009a-c Addition of carbon tetrachloride to unsaturated hydrocarbons TIAsscher, Meir; Vofsi, David; Katchalsky, Aharon IN SO 17 pp. DTPatent LΑ Unavailable KIND DATE PATENT NO. APPLICATION NO. DATE ______ FR 1334749 PΙ 19630809 19610928 PRAI IL Styrene (10.4 g.) and 31 g. CCl4 was heated 15 hrs. at 100° with AB 0.17 g. CuCl2.2H2O and 0.6 g. Et2NH2Cl in 15 g. MeCN to give 84% 1,1,1,3-tetrachloro-3-phenylpropane. Benzoin (106 mg.) in 15.4 ml. CCl4 and 135 mg. FeCl3.6H2O and 82 mg. Et2NH.HCl in 10 g. tert-BuOH was treated in an air-free autoclave with 22 g. ethylene at 90° for 16 hrs. to give 39% 1,1,1,3-tetrachloropropane, 50% 1,1,1,5-tetrachloropentane, and 11% 1,1,1,7-tetrachloro-n-heptane. In the absence of benzoin, only trace quantities of the tetrachloro compds. were formed. The following RCH2CCl3 were prepared (R, b.p., and n25D given): PhCHCl, b0.1 84°, 1.5523;

PhCHOMe, b0.5 80°, 1.5280; ClCH2CH:CH, b25 110°, 1.5042; MeOCH2CH:CH, b20 103°, 1.4792; NCCHCl, b25 111°, 1.4931; MeO2CCHCl, b20 111°, 1.4819; HOCH2CHCl, b1.0 66°, 1.5066; EtCHCl, b25 87°, 1.4779; C6H13CHCl, b0.8 70°, 1.4749.

MeCHClCHMeCCl3, b25 93°, n25D 1.4869, was also prepared

L5 ANSWER 33 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN

AN 1963:461505 CAPLUS

DN 59:61505

OREF 59:11248d-h

TI Addition compounds of alkenes with chloroform

PA Meir Asscher, Aharon Katchalsky, and David Vofsi

SO 12 pp.

DT Patent

LA Unavailable

PATENT NO. KIND DATE APPLICATION NO. DATE
BE 622938 19630115 BE

PI BE 622938 19 PRAI IL 19

19610928 CHCl2CHRCHZR1 (I) and CHCl2CHRCR1: CR2CHZR3 (II) [R, R1, R2, R3, H, alkyl, aryl, arylalkyl; Z, Cl, alkoxyl, useful intermediates for the preparation of perfumes, were prepared by reaction of CHCl3 with an alkene, arylalkene, or alkanediene in the presence of halogen transferring agents (as Cu or Fe salts in the one or other state of their valency) under addition of an amine hydrochloride to give excess Cl ions in ratio to the metal ions; the process is improved by addition of a reducing agent (e.g. benzoin, a sulfite) to hold at least a part of the halogen transferring agent in its lower valence during the reaction. Thus, 11.2 g. 1-octene, 36 g. CHCl3, 0.5 g. FeCl3.6H2O, 0.5 g. Me2NH.HCl, and 15 g. MeOH heated 22 hrs. at 142° in a closed apparatus, cooled, the mixture washed with N HCl and H2O, and the solution distilled gave 80% I (R = H, R1 = n-C6H13, Z = Cl), b10 71-81°. Butadiene (8.1 g.) was dissolved in a solution of 0.318 g. benzoin in 54 g. CHCl3, a solution of 0.405 g. FeCl3.6H2O and 0.246 g. Et2NH.HCl in 10 g. MeCN added, the mixture heated 16 hrs. at 130° in a Carius tube in which the air is replaced by butadiene, cooled (ice-salt), washed (0.1N HCl), and distilled to give 9.3 g. of a fraction, b25 100-40°, containing 90%II(R = R1 = R2 = R3 = H, Z = C1) and 6.8 g. of a fraction, b0.0606 85-135°, containing a compound C9H13Cl3. Similarly, optionally in the presence of benzoin and (or) Et2NH.HCl, the following I were prepared (R, Et, Cl, 75°/25, 1.4619; Me, Me, Cl, 76°/25, 1.4683; H, Ph, Cl, 62°/0.1, --; H, Ph, MeO, 76°/0.3, 1.5192. Also the following II were prepared (R, R1, R2, R3, Z, b.p./mm., and n25D given): H, H, H, H, MeO, 105°/20, 1.4792; H, H, H, H, Cl, 100°/25, 1.4969. Belg. 622,939; 17 pp. Addition compds. of CCl4 with substituted hydrocarbons (containing at least one nonaromatic double bond or a pair of conjugated double bonds) were prepared to give useful intermediates for the preparation of insecticides and pharmaceuticals; in the case of rubber a highly chlorinated product was obtained. The following compds. were prepared (product, b.p./mm., and n25D given): PhCHClCH2CCl3, 84°/0.1, 1.5523; PhCH(OMe)CH2CCl3, 80°/1.5, 1.5280; CCl3CH2CH:CHCH2Cl, 110°/25, 1.5042 (by-product was tetrachlorononadiene, b0.08 50-130°); CCl3CH2CH:CHCH2OMe, 103°/20, 1.4792; CCl3CH2CHClCN, 111°/25, 1.4931; CCl3CH2CHClCO2Me, 111°/20, 1.4819; CCl3CH2CHClCH2OH, 66°/1.0, 1.5066; a mixture (prepared from H2C:CH2) containing CH2ClCH3CCl3, CH2Cl(CH2)3CCl3, and CH2Cl(CH2)5CCl3, in 39:50:11 weight-% ratio, 60-152°/20, --; CCl3CH2CHClEt, 87°/25, 1.4779; MeCHClCH(CCl3)Me, 93°/25, 1.4869; CCl3CH2CHClC6H13, 70°/0.8, 1.4749; a brown-gray powder containing 49.54% Cl and prepared from a solution containing 5% latex of natural rubber, --, --; Me(CH2)5CHClCH2CCl3, 87-95°/0.3, 1.4746.

L5 ANSWER 34 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN

AN 1963:59294 CAPLUS

DN 58:59294

OREF 58:10063d-e

PΤ

AB

IL 13845

GB 920855

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TI
     Chlorine activation by redox transfer. II. The addition of carbon
     tetrachloride to olefins
ΑU
     Asscher, M.; Vofsi, D.
CS
     Weizmann Inst. Sci., Rehovoth, Israel
SO
     Journal of the Chemical Society, Abstracts (1963) 1887-96
     CODEN: JCSAAZ; ISSN: 0590-9791
DT
     Journal
     Unavailable
LA
     cf. CA 55, 24539d. CCl4 is added to olefins and vinylic monomers under
AB
     the catalytic influence of iron (II, III) or copper (I, II)
     chloride, in a variety of solvents at 70-145°. Copper chloride
     completely suppresses telomerization. A free-radical chain mechanism is
     proposed for the addition in which metal chloride participates in the
     propagation as a chlorine atom transfer agent, thereby much enhancing the
     apparent reactivity of CCl4 ("redox-transfer"). Exptl. support for this
     mechanism is presented. The reaction with but-2-ene gives a mixture of 1:1
     diastereoisomeric adducts, the composition of which changes with catalyst, and,
     for copper catalysis, also with the solvent and with the excess of
     chloride ion. cis- and trans-But-2-ene give the same ratio of
     diastereoisomers. The initiation mechanism is discussed. At 82°,
     ferric chloride induces addition of CCl4 to but-2-ene only in the
     presence of a reducing agent. This enables the estimation of the kinetic chain
     length, which has a much larger value than for a peroxide-induced
     reaction.
    ANSWER 35 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
L_5
    1963:461508 CAPLUS
DN
    59:61508
OREF 59:11249c-d
ΤI
    Telomers
IN
    Takahashi, Akira; Mogi, Noboru; Takahama, Hiroshi
PA
     Showa Denko K. K.
SO
    7 pp.
    Patent
DT
    Unavailable
T.A
    PATENT NO. KIND DATE
                                        APPLICATION NO. DATE
                                         -----
                                                          -----
    JP 37018389 19621128
US 3213149 1965
PI
                                         JΡ
                                                          19610301
                                         US
AB
    Telomers were prepared by the reaction of ethylene or vinyl compds. and
     telogens in the presence of amines and heavy metal salts of organic acids.
     Thus, 476 parts CCl4, 1.65 parts triethanolamine, and iron
    naphthenate (containing 0.034 part Fe) in an autoclave under N was kept at 50
     atmospheric ethylene for 4 hrs. at 70°, cooled, ethylene and CCl4 evaporated,
     and the telomer obtained in 214 parts yield; it gave 5 fractions of
     Cl(CH2CH2)nCCl3 (% yield, n, b.p. given): 8.2, 1, b24 58-61°; 53.2,
     2, b24 111-13°; 22.3, 3, b2 92-4°; 10.3, 4, b2
    120-4°; 6.0, ≥5, --).
L_5
    ANSWER 36 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN
AN
    1962:52908 CAPLUS
DN
    56:52908
OREF 56:9961a-c
TI
    Telomers or adducts
IN
    Katchalsky, Aharon; Vofsi, David; Asscher, Meir; Levy, Edmond
DT
    Patent
LA
    Unavailable
    PATENT NO.
                                         APPLICATION NO. DATE
                    KIND DATE
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                                         ______
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19611123

Telomers formed by the reaction between CC14 as telogen and an

IL

GB

 α -olefinic compound as taxogen may be prepared using an organic amine or ammonia as primary catalyst and cupric or ferrous salts as secondary catalyst. A sequestering agent such as ethylenediaminetetraacetic acid may also be added. The telomerization may be performed either in aqueous or in anhydrous medium. Safety hazards involved in the use of conventional type catalysts (benzoyl peroxide, azo compds.) are thus eliminated. The 12 examples given include the preparation of Cl(CH2CH2)nCCl3 (n = 1 through 5) and 1,1,1,3-tetrachloro-n-nonane.

L5 ANSWER 37 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN

AN 1955:23480 CAPLUS

DN 49:23480

OREF 49:4496f-i,4497a-c

Chemical transformations of the trichloromethyl group in saturated polychlorohydrocarbons

ΑU Nesmeyanov, A. N.; Freidlina, R. Kh.; Zakharkin, L. I.

SO Doklady Akademii Nauk SSSR (1954), 96, 87-90 CODEN: DANKAS; ISSN: 0002-3264

DTJournal LAUnavailable cf. C.A. 47, 3789c. The CCl3 group in polychloroalkanes is inert toward AΒ nucleophilic agents. 1,1,1-Trichloropentane does not react with NH3 in EtOH even at 160° or liquid NH3 at 140°, with NaI, or NaHC(CO2Et)2. Other terminal trichloroalkanes give the same result. $\alpha, \alpha, \alpha, \omega$ -tetrachloroalkanes only the terminal lone Cl reacts with nucleophilic agents. Thus NaI in Me2CO reacts with 1,1,1,5-tetrachloropentane yielding 90% 1,1,1-trichloro-5-iodo-pentane, b1.5 78-9°, nD20 1.5480, d20 1.8086; the product is readily converted to the cyano derivative by treatment with KCN. Reaction of KOAc with 1,1,1,5-tetrachloropentane in hot AcOH 18 h. in presence of a little KI gave 86% 1,1,1-trichloro-5-acetoxypentane, b3.5 99-100°, nD20 1.4700, d20 1.2859; this was readily converted to 5,5,5-trichloro-1pentanol, b10 112-12.5°, nD20 1.4897, d20 1.3431; other nucleophilic agents give similar results. Electrophilic agents attack the CCl3 group preferentially (cf. C.A. 49, 2298b); the group being resistant to hydrolysis in basic solution, is rapidly attacked by acid solns. and in concentrated H2SO4 readily yields the CO2H group. While CC13 group does not display Cl exchange for Br in presence of HBr it does so in the presence of electrophilic AlCl3. Passage of HBr into 1,1,1-trichloropentane at 4-5° in the presence of a little AlCl3 gave a good yield of 1,1,1-tribromopentane, b8 85-6°, nD20 1.5390, d20 1.9882. Similarly 1,1,1,5-tetrachloropentane yields 1,1,1-tribromo-5chloropentane, b2 101-2°, nD20 1.5655, d20 2.0902. The cleavage of HCl from 1,1,1,2-tetrachloropentane by H2SO4 begins only at 140-60°; similar behavior is seen in 1,1,1,2,5-pentachloropentane. The action of FeCl3 leads to HCl loss from 1,1,1,5-tetrachloropentane at 30-50°; 1,1,1,2,5-pentachloropentane requires 100-20°, yielding 1,1,2,5-tetrachloro-1-pentene, b8 92-3°, nD20 1.5113, d20 1.4121. Addition of Cl with cooling to 1,1-dichloro-1-pentene and 1,1,5-trichloro-1-pentene gave, resp., 1,1,1,2-tetrachloropentane, b8 72-3°, nD20 1.4825, d20 1.3339, and 1,1,1,2,5-pentachloropentane, b12 121-2°, nD20 1.5135, d20 1.4807. Action of free radical type reagents was examined in the action of 1,1,1,5-tetrachloropentane with PhMgBr in the presence of CoCl2, with Raney Ni and powdered Cu. In all cases the reaction took place at the CCl3 group. No reaction took place with PhMgBr alone; in the presence of CoCl2 there was formed a mixture of products from which were isolated Ph2 and 1,5,5,6,6,10-hexachlorodecane (I); the products evidently arise from radical attack and the latter is accounted for by formation of Cl(CH2)4CCl2 radical. Action of Raney Ni in EtOH gave, after 2 h. reflux, 37% I; powdered Cu gave the same result. While heterolytic reactions in terminal CH2Cl group by nucleophilic agents, and in CCl3 group by electrophilic agents proceeds well and give few

10/648,709

byproducts, the homolytic reactions of CCl3 group are apt to give product mixture. The higher polychloroalkanes behave in the same way as the above-described pentanes. However 1,1,1,3-tetrachloropropane reacts differently; with NaCN, Na2S or other nucleophilic agents it does not exchange Cl in the CH2Cl group but undergoes HCl cleavage forming the olefin derivs.

L5 ANSWER 38 OF 38 CAPLUS COPYRIGHT 2004 ACS on STN

AN 1951:3629 CAPLUS

DN 45:3629

OREF 45:635d-f

TI 1,1,1,6,6,6-Hexachlorohexane

IN Neuworth, Martin B.

PA Socony-Vacuum Oil Co., Inc.

DT Patent

LA Unavailable

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI US 2519691 19500822 US

AB CCl3(CH2)2Cl (I) heated with Fe, Cu, or Al at 100-150° in the presence of H2O yields CCl3(CH2)4CCl3 (II), m. 108-9° (from EtOH). Thus 30 cc. I, 25 g. powdered Fe, and 150 cc. H2O are refluxed with stirring for 28 hrs., the mixture extracted with Et2O, and the Et2O evaporated to leave an

oily residue from which II crystallizes in colorless needles (20% yield). II is useful as plasticizer for chlorinated resins and as intermediate for adipic acid.

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